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- THE LUBRIZOL CORPORATION (71) Applicant: [US/US]; 29400 Lakeland Blvd., Wickliffe, Ohio 44092-2298 (US).
- (72) Inventors: BURRINGTON, James D.; 6858 Bramblewood Lane, Mayfield Village, Ohio 44143 (US). DAILY, Daniel T.; 33166 Popham Lane, Solon, Ohio 44139 (US). GEORGE, Herman F.; 11840 Clarkwood Drive, Chardon, Ohio 44024 (US).
- GILBERT, TERESAN W.; ESPOSITO, (74) Agent: MICHAEL F.; LAFERTY, DAVID M.; The Lubrizol Corporation, Patent Dept./022B, 29400 Lakeland Blvd., Wickliffe, Ohio 44092-2298 (US).

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(54) Title: GEL ADDITIVES FOR FUEL THAT REDUCE SOOT AND/OR EMISSIONS FROM ENGINES

(57) Abstract: An additive gel for fuels that reduces the soot content in lubricating oil and/or emission in an engine. Further a process employing an additive gel for fuel in a fuel system to decrease the amount of soot in the lubricating oil of an engine and/or decrease the emissions from an engine.

# Title: GEL ADDITIVES FOR FUEL THAT REDUCE SOOT AND/OR EMISSIONS FROM ENGINES

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#### **Background of the Invention**

The present invention relates to a novel gel composition that is an additive for fuel that results in a decrease in the amount of soot in a lubricating oil in an engine and/or a decrease in the amount of emissions particularly soot, hydrocarbons and/or nitrogen oxides, (NO, NO<sub>2</sub>, N<sub>2</sub>O collectively NOx) from an engine.

A complication facing modern compression ignited and spark ignited engines is the build up of soot in the lubricating oil due to oxidation and nitration by-products of the unburnt fuel or the lubricating oil itself and environmentally unfriendly emission. The buildup of this soot thickens the lubricating oil and can cause engine deposits. In severe operating conditions, the oil can thicken to the point of gelling. When the soot levels get to high, the increase in oil viscosity results in poor lubrication at critical wear points on the engine. This poor lubrication results in high wear, the formulation of higher amounts of piston deposits, a loss in fuel economy occurs and increased exhaust emissions. The net result is a shorter effective life of the lubricating oil and exhaust emissions.

Another complication in facing modern and future engines is the need for these engines to meet upcoming emission legislation. One solution has been to use exhaust after treatment systems to reduce emissions for the engine.

It is desirable to decrease the concentration of particles of soot in an engine oil using a novel gel composition as an additive to fuel. It is further desirable to decrease the emissions of soot, hydrocarbons and/or NOx from an engine using a novel gel fuel additive.

It has been found that a gel fuel additive in contact with the fuel of an engine can decrease the soot content in the oil of the lubricating system of the engine. It has been further found that an gel fuel additive can reduce the emissions from an engine in particular soot, hydrocarbons and/or NOx emissions.

This invention provides a way to provide enhanced performance to lubricating oil and to reduce engine emissions into the environment.

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### **Summary of the Invention**

In accordance with the instant invention, it has been discovered that a gel additive for fuel can reduce the concentration of soot particles in a lubricating oil and/or reduce emissions from an engine.

In accordance with the present invention it has been discovered that a gel additive for fuel comprising a dispersant and a detergent ("gel") reduces the concentration of soot in the lubricating oil of an engine and/or decreases the emissions from an engine. Further the gel can contain an antioxidant and also may contain other fuel soluble additives. The gel additive for fuel is fuel soluble. The gel dissolves during use of the engine. In one embodiment the release of the gel components is a slow release.

In the present invention, suspended and/or dispersed soot in engine oil is decreased by a process comprising contacting a portion of the fuel in an engine with a gel. Further the present invention decreases the emissions and soot, hydrocarbons and/or NOx from an engine by a process comprising contacting a portion of the fuel of an engine with a gel.

The present invention provides for the use of a gel to decrease the amount of suspended/dispersed soot in lubricating oil in engines and/or to decrease the emissions in particular soot, hydrocarbons and/or NOx from an engine. The engines that can use the gel include, but are not limited to internal combustion engines such as spark ignited and/or a compression ignited, stationary and/or mobil power plant engines, generators, diesel and/or gasoline engines, on highway and/or off highway engines, two-cycle engines, aviation engines, piston engines, marine engines, railroad engines, biodegradable fuel engines and the like. In one embodiment the engine is equipped with after treatment devices, such as exhaust gas recirculation systems, catalytic converters, diesel particulate filters, NOx traps and the like.

### **Detailed Description of the Invention**

In accordance with the present invention the soot concentration is decreased from a lubricating oil in an engine thereby avoiding the deleterious effects on the

engine from the soot, including viscosity and wear. Furthermore, the emissions of an engine is decreased thereby improving the environment.

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The soot level in the lubricating oil and/or in the emissions from the engine is reduced by contacting the fuel with the gel. The gel is positioned within the fuel system, anywhere the gel will be in contact with the fuel. The gel is positioned anywhere that the circulating fuel contacts the gel such as full flow of fuel, bypass of the fuel or combinations therein. The location of the gel in the fuel system includes but is not limited to a filter, fuel filter, fuel bypass loop, fuel pump, injectors canister, housing, reservoir, pockets of a filter, canister in a filter, mesh in a filter, canister in a bypass system, mesh in a bypass system, canister in a tank, mesh in a tank, manifolds, inlets and/or outlets of fuel tank, fill fuel pipe, valves in fuel system, fuel chambers, fuel drain, intake air system, positive crank case ventilation system, air intake filter, exhaust gas recirculation (egr) system and the like. One or more locations can contain the gel. Further, if more than one location is used the gel can be identical, similar and/or a different formulation.

A necessary design feature for the application of the gel is that a portion to all of the gel components reach the combustion chamber. In one embodiment it is desirable to provide a container to hold the gel, such as a housing, a canister, a structural mesh or the like anywhere within the fuel system, for example, a housing in the filter of the fuel system. In one embodiment the design feature for the container is that at least a portion of the gel is in contact with the fuel and/or the components of the gel reach the combustion chamber.

In one embodiment, the gel is positioned anywhere in the fuel filter. The fuel filter is a desirable location to place the gel because the gel and/or spent gel can easily be removed, and then replaced with a new and/or recycled gel. In another embodiment the gel is positioned in a container anywhere in the fuel tank.

The gel or a portion of the gel needs to be in contact with the fuel. In one embodiment the gel is in contact with the fuel in the range of about 100% to about 1% of the fuel, in another embodiment the gel is in contact with the fuel in the range of about 75% to about 25% of the fuel and in another embodiment the gel is in contact with the fuel in the range of about 50% of the fuel.

The gel can be added also to the fuel by the fuel supplier at a refinery, terminal, or at a refueling station by premixing the gel with the fuel. Alternatively, the vehicle operator can add the gel to the fuel tank by dosing the tank during refueling. The gel additive may be dosed to the fuel using a fuel dosing system that provides a controlled level of the additive to the fuel (storage) tank.

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The release rate of the gel is determined primarily by the gel formulation. Also the location and the flow rate of the fuel and/or air affects the rate at which the gel dissolves. In one embodiment the gel is positioned in a location of a high flow rate such as about 50% to about 100% of the circulating fuel. In another embodiment the gel is positioned in a location of medium flow rate such as about 25% to about 75% of the circulating fuel. In another embodiment the gel is positioned in a location of low flow rate such as ≥1% to about 25% of the circulating fuel. For a given gel formulation the flow rate of the circulating fuel is directly proportional to the dissolution rate of the gel. Therefore as the flow rate decreases there is less dissolution of the gel and as the flow rate increases there is greater dissolution of the gel. The gel is positioned in a location desirable for the specified and desirable dissolution rate of the gel.

In one embodiment the gel's formulation may be composed of one or more components such as fuel soluble additives so that at the end of its service life there is none to little gel residue remaining. In another embodiment the gel's formulation maybe composed one or more component that selectively dissolve while at least a portion of the components remain at the end of its service life.

The gel for the fuel system comprises a dispersant, a detergent, and an antioxidant. Further the gel may optionally contain other fuel soluble additives.

In one embodiment the gel is represented by the formula A+B+C wherein A equals at least one component with at least one or more reactive or associative groups; wherein B contains a particle(s) or other component(s) with at least one group which reacts or associates with A to form a gel, and wherein C is at least one or more desired fuel additives. In one embodiment the gel has an antioxidant, a detergent and dispersant.

Component A includes but is not limited to antioxidants; dispersants; ashless dispersants; succinics; maleic anhydride styrene copolymers; maleated ethylene diene monomer copolymers; surfactants; emulsifiers; functionalized derivatives of each component listed herein and the like; and combinations thereof. Component A can be used alone or in combination. In one embodiment the preferred A is polyisobutenyl succinimide dispersant.

Component B includes but is not limited to dispersants, detergents, overbased detergents, carbon black, silica, alumina, titania, magnesium oxide, calcium carbonate, lime, clay, zeolites and the like; and combinations thereof. Component B can be used alone or in combination. In one embodiment Compound B is an overbased alkybenzenesulfonate detergent.

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Component C includes but is not limited to the additives which include but are not limited to antioxidant, extreme pressure (EP) agents, wear reduction agents, viscosity index improvers, anti-foaming agents, combustion modifiers, cetane improver, fuel dispersants and the like. Component C can be used alone or in combination. In one embodiment Component C is at least one of an antioxidant and if component A is an antioxidant they are not the same antioxidant. In one embodiment Component C is at least one of a dispersant and if component A is a dispersant they are not the same dispersant.

The gel contains component A in the range of about 0.1 % to about 95 %, in one embodiment about 1% to about 70% and in another embodiment about 7% to about 50% of the gel. The gel contains component B in the range of about 0.1% and about 99%, in one embodiment about 5% to about 80% and in another embodiment about 10% to about 70% of the gel. The gel contains component C in the range of about 0% to about 95%, and in one embodiment about 1% to about 70% and in another embodiment about 5% to about 60% of the gel.

In accordance with the present invention the gel formed is a fuel based gel.

The gel is selected from the group comprising at least one of dispersants, dispersant precursors (such as alkyl or polymer succinic anhydrides), detergents, antioxidants, and mixtures thereof. Optionally, at least one soluble additive may be added to the gel as desired.

Other fuel soluble additives include, but are not limited to friction reducing agents, extreme pressure (EP) agents, wear reduction agents, viscosity index improvers, anti-foaming agents, anti-misting agents, cloud-point depressants, pourpoint depressants, mineral or synthetic oils, anti-knock agents, lead scavengers, dyes, cetane improvers, rust inhibitors, bacteriostatic agents, gum inhibitors, fluidizers, metal deactivators, demulsifiers, anti-icing agents, lubricity additives, friction modifiers, viscosity improvers, flow improvers, low temperature improvers, anti-static agents, valve-seat recession agents, intake valve deposit control additives, combustion chamber deposit control additives, fuel injector deposit control additives and the like. The fuel soluble additives may be used alone or in combination. The gel contains the fuel soluble additives in the range of about 0 % to about 90 %, and in one embodiment about .0005 % to about 50 % and in another embodiment about .0025 % to about 30% of the gel.

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Optionally, the fuel can also contain fuel-borne catalysts (organometallic compounds of e.g. Na, K, Co, Ni, Fe, Cu, Mn, Mo, Va, Zi, Be, Pt, Pa, Ce, Cr, Al, Th, Se, Bi, Cd, Te, Th, Sn, Ba, B, La, Ta, Ti, W, Zn, Ga, Pb, Ag, Au, Os, Ir) mixtures thereof and the like.

The gel typically contains small amounts (about 5-40%) of a hydrocarbon base, which include but are not limited to petroleum-based fuels, synthetics or mixtures thereof.

The gel comprises mixtures of two or more substances and exists in a semi-solid state more like a solid than a liquid. The rheological properties of a gel can be measured by small amplitude oscillatory shear testing. This technique measures the structural character of the gel and produces a term called the storage modulus (which represents storage of elastic energy) and the loss modulus (which represents the viscous dissipation of that energy). The ratio of the loss modulus/storage modulus, which is called the loss tangent, or "tan delta," is >1 for materials that are liquid-like and <1 for materials that are solid-like. The gels have tan delta values in one embodiment of about  $\leq 0.75$ , in one embodiment of about  $\leq 0.5$  and in one embodiment of about  $\leq 0.3$ .

In one embodiment the gels are those in which gelation occurs through the combination of a detergent and a dispersant in particular on overbased detergent and ashless succimide dispersed. In this embodiment, the ratio of the detergent to the dispersant is typically from about 10:1 to about 1:10; in one embodiment from about 5:1 to about 1:5; in one embodiment from about 4:1 to about 1:1; and in one embodiment from about 4:1 to about 2:1. In addition, the TBN (total base number) of the overbased detergents is in one embodiment at least 100, in one embodiment at least 300, in one embodiment at least 400 and in one embodiment 600. Where mixtures of overbased detergents are used, at least one should have a TBN value of at least 100. However, the average TBN of these mixtures may also correspond to a value greater than 100.

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The dispersants include but are not limited to ashless-type dispersants, polymeric dispersants, Mannich dispersants, high molecular weight (Cn wherein n ≥12) esters, carboxylic dispersants, amine dispersants, amine dispersants, polymeric dispersants and combinations thereof. The dispersant may be used alone or in combination. The dispersant is present in the range from about 0.1% to about 95% of the gel, preferably from about 1% to about 70% of the gel, and preferably from about 7% to about 50% of the gel.

The dispersant in the gel includes but is not limited to an ashless dispersant such as a polyisobutenyl succinimide and the like. Polyisobutenyl succinimide ashless dispersants are commercially-available products which are typically made by reacting together polyisobutylene having a number average molecular weight ("Mn") of about 300 to 10,000 with maleic anhydride to form polyisobutenyl succinic anhydride ("PIBSA") and then reacting the product so obtained with a polyamine typically containing 1 to 10 ethylene diamine groups per molecule.

Ashless type dispersants are characterized by a polar group attached to a relatively high molecular weight hydrocarbon chain. Typical ashless dispersants include N-substituted long chain alkenyl succinimides, having a variety of chemical structures including typically:

$$R_1$$
 $R_1$ 
 $R_1$ 
 $R_2$ 
 $R_2$ 
 $R_2$ 
 $R_2$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_4$ 
 $R_4$ 
 $R_4$ 

and/or

$$R_1$$
 $N$ -[ $R_2$ NH]x- $R_2$ N

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wherein each R<sup>1</sup> is independently an alkyl group, frequently a polyisobutyl group with a molecular weight of 500-5000, and R<sup>2</sup> are alkenyl groups, commonly ethylenyl (C<sub>2</sub>H<sub>4</sub>) groups. Succinimide dispersants are more fully described in U.S. Patent 4,234,435 which is incorporated herein by reference. The dispersants described in this patent are particularly effective for producing gels in accordance with the present invention.

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The Mannich dispersant are the reaction products of alkyl phenols in which the alkyl group contains at least about 30 carbon atoms with aldehydes (especially formaldehyde) and amines (especially polyalkylene polyamines). Mannich bases having the following general structure (including a variety of different isomers and the like) are especially interesting.

and/or

$$H_2C$$
 $H_2C$ 
 $H_2C$ 

Another class of dispersants is carboxylic dispersants. Examples of these "carboxylic dispersants" are described in Patent 3,219,666.

Amine dispersants are reaction products of relatively high molecular weight aliphatic halides and amines, preferably polyalkylene polyamines. Examples thereof are described, in U.S. Patent 3,565,804.

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Polymeric dispersants are interpolymers of oil-solubilizing monomers such as decyl methacrylate, vinyl decyl ether and high molecular weight olefins with monomers containing polar substituents, e.g., aminoalkyl acrylates or acrylamides and poly-(oxyethylene)-substituted acrylates. Examples of polymer dispersants thereof are disclosed in the following U.S. Patents: 3,329,658, and 3,702,300.

Dispersants can also be post-treated by reaction with any of a variety of agents. Among these are urea, thiourea, dimercaptothiadiazoles, carbon disulfide, aldehydes, ketones, carboxylic acids, hydrocarbon-substituted succinic anhydrides, nitriles, epoxides, boron compounds, and phosphorus compounds.

The detergents include but are not limited to overbased sulfonates, phenates, salicylates, carboxylates and the like, overbased calcium sulfonate detergents which are commercially-available, overbased detergents containing metals such as Mg, Ba, Sr, Na, Ca and K and mixtures thereof. The detergents may be used alone or in combination. Detergents are described, for example, in U.S. Patent 5,484,542 which is incorporated herein by reference. The detergents are present in the range from about 0.1% to about 99%, preferably from about 5% to about 80% and more preferably from about 10% to about 70% by weight of the gel.

Antioxidants include but are not limited to alkyl-substituted phenols such as 2, 6-di-tertiary butyl-4-methyl phenol, phenate sulfides, phosphosulfurized terpenes, sulfurized esters, aromatic amines, diphenyl amines, alkylated diphenyl amines and hindered phenols.

The antioxidant includes amine antioxidants and is not limited to bisnonylated diphenylamine, nonyl diphenylamine, octyl diphenylamine, bis-octylated diphenylamine, bis-decylated diphenylamine, decyl diphenylamine and mixtures thereof.

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The antioxidant includes sterically hindered phenols and includes but is not limited to 2,6-di-tert-butylphenol, 4-methyl-2,6-di-tert-butylphenol, 4-ethyl-2,6-di-tert-butylphenol, 4-propyl-2,6-di-tert-butylphenol, 4-butyl-2,6-di-tert-butylphenol 2,6-di-tert-butylphenol, 4-pentyl-2,6-di-tert-butylphenol, 4-hexyl-2,6-di-tert-butylphenol, 4-hexyl-2,6-di-tert-butylphenol, 4-(2-ethylhexyl)-2,6-di-tert-butylphenol, 4-octyl-2,6-di-tert-butylphenol, 4-nonyl-2,6-di-tert-butylphenol, 4-dodecyl-2,6-di-tert-butylphenol, 4-dodecyl-2,6-di-tert-butylphenol, 4-tridecyl-2,6-di-tert-butylphenol, 4-tetradecyl-2,6-di-tert-butylphenol, methylene-bridged sterically hindered phenols include but are not limited to 4,4'-methylenebis(6-tert-butyl-0-cresol), 4,4'-methylenebis(2-tert-amyl-o-cresol), 2,2'-methylenebis(4-methyl-6-tert-butylphenol), 4,4'-methylene-bis(2,6-di-tertbutylphenol) and mixtures thereof.

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Another example of an antioxidant is a hindered, ester-substituted phenol, which can be prepared by heating a 2,6-dialkylphenol with an acrylate ester under base catalysis conditions, such as aqueous KOH. Antioxidants may be used alone or in combination.

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The antioxidants are typically present in the range of about 0.01% to about 95%, preferably about 0.01% to 95%, and more preferably about 1.0% to about 70% and most preferably about 5% to about 60% by weight of the gel.

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The extreme pressure anti-wear additives include but are not limited to a sulfur or chlorosulphur EP agent, a chlorinated hydrocarbon EP agent, or a phosphorus EP agent, or mixtures thereof. Examples of such EP agents are chlorinated wax, organic sulfides and polysulfides, such as benzyldisulfide, bis-(chlorobenzyl) disulfide, dibutyl tetrasulfide, sulfurized sperm oil, sulfurized methyl ester of oleic acid sulfurized alkylphenol, sulfurized dispentene, sulfurized terpene, and sulfurized Diels-Alder adducts; phosphosulfurized hydrocarbons, such as the reaction product of phosphorus sulfide with turpentine or methyl oleate, phosphorus

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esters such as the dihydrocarbon and trihydrocarbon phosphate, *i.e.*, dibutyl phosphate, diheptyl phosphate, dicyclohexyl phosphate, pentylphenyl phosphate; dipentylphenyl phosphate, tridecyl phosphate, distearyl phosphate and polypropylene substituted phenol phosphate, metal thiocarbamates, such as zinc

dioctyldithiocarbamate and barium heptylphenol diacid, such as zinc dicyclohexyl phosphorodithioate and the zinc salts of a phosphorodithioic acid combination may be used and mixtures thereof. The EP agent can be used alone or in combination. The EP agents are present in the range of about 0% to 10%, preferably from about 0.25% to about 5% and more preferably from about 0.5% to about 2.5% by weight of the gel.

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The viscosity modifiers provide both viscosity improving properties and dispersant properties. Examples of dispersant-viscosity modifiers include but are not limited to vinyl pyridine, N-vinyl pyrrolidone and N,N'-dimethylaminoethyl methacrylate are examples of nitrogen-containing monomers and the like. Polyacrylates obtained from the polymerization or copolymerization of one or more alkyl acrylates also are useful as viscosity modifiers. The viscosity modifiers may be used alone or in combination.

Functionalized polymers can also be used as viscosity modifiers. Among the common classes of such polymers are olefin copolymers and acrylate or methacrylate copolymers. Functionalized olefin copolymers can be, for instance, interpolymers of ethylene and propylene which are grafted with an active monomer such as maleic anhydride and then derivatized with an alcohol or an amine. Other such copolymers are copolymers of ethylene and propylene which are reacted or grafted with nitrogen compounds. Derivatives of polyacrylate esters are well known as dispersant viscosity index modifiers additives. Dispersant acrylate or polymethacrylate viscosity modifiers such as Acryloid<sup>TM</sup> 985 or Viscoplex<sup>TM</sup> 6-054, from RohMax, are particularly useful. Solid, oil-soluble polymers such as the PIB, methacrylate, polyalkylstyrene, ethylene/propylene and ethylene/propylene/1, 4-hexadiene polymers, can also be used as viscosity index improvers. The viscosity modifiers are known and commercially available. The viscosity modifiers are

present in the ranged about 0% to about 20%, preferably about 5% to about 15% and more preferably about 7% to about 10% of the gel.

The antifoams include but are not limited to organic silicones such as poly dimethyl siloxane, poly ethyl siloxane, poly diethyl siloxane and the like. The antifoams may be used alone or in combination. The antifoams are normally used in the range of about 0% to about 1%, preferably about 0.02% to about 0.5% and more preferably 0.05% to about 0.2% by weight of the gel.

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The anti knock agents, includes but is not limited to tetra-alkyl lead compounds, organomanganese compounds and the like. The anti knock agents may be used alone or in combination. The anti knock agents are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30 % of the gel.

The lead scavengers, includes but is not limited to halo-alkanes and the like. The lead scavengers may be used alone or in combination. The lead scavengers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The dyes, includes but is not limited to halo-alkanes and the like. The dyes may be used alone or in combination. The dyes are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The combustion modifiers, includes but is not limited to alkyl nitro compounds and the like. The combustion modifiers may be used alone or in combination. The combustion modifiers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The cetane improvers, includes but is not limited to alkyl nitrates and the like. The cetane improvers may be used alone or in combination. The cetane improvers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The rust inhibitors, includes but is not limited to alkylated succinic acids and anhydrides derivatives thereof, organo phosphonates and the like. The rust inhibitors may be used alone or in combination. The rust inhibitors are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

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The bacterostatic agents, includes but is not limited to formaldehyde, gluteraldehyde and derivatives, kathan and the like. The bacterostatic agents may be used alone or in combination. The bacterostatic agents are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.78

The gum inhibitors, includes but is not limited to diphenyl amine and the like. The gum inhibitors may be used alone or in combination. The gum inhibitors are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The fluidizers, includes but is not limited to polyisobutenyl amine, polypropylene oxide, and the like. The fluidizers may be used alone or in combination. The fluidizers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The metal deactivators, includes but is not limited to derivatives of benzotriazoles such as tolyltriazole, N,N-bis(heptyl)-ar-methyl-1H-benzotriazole-1-methanamine, N,N-bis(nonyl)-ar-methyl-1H-Benzotriazole-1-methanamine, N,N-bis(decyl)-ar-methyl-1H-Benzotriazole-1-methanamine, N,N-bis(undecyl)-ar-methyl-1H-benzotriazole-1-methanamine, N,N-bis(dodecyl)-ar-methyl-1H-Bbenzotriazole-1-methanamine N,N-bis(2-ethylhexyl)-ar-methyl-1H-Bbenzotriazole-1-methanamine and mixtures thereof. In one embodiment the metal deactivator is N,N-bis(2-ethylhexyl)-ar-methyl-1H-benzotriazole-1-methanamine; 1,2,4-triazoles, benzimidazoles, 2-alkyldithiobenzimidazoles; 2-alkyldithiobenzothiazoles; 2-(N,N-dialkyldithiocarbamoyl)benzothiazoles; 2,5-bis(alkyl-dithio)-1,3,4-thiadiazoles such as 2,5-bis(tert-octyldithio)-1,3,4-

thiadiazole 2,5-bis(tert-nonyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-decyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-dodecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-dodecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-tetradecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-pentadecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-pentadecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-hexadecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-hexadecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-octadecyldithio)-1,3,4-thiadiazole, 2,5-bis(tert-eicosyldithio)-1,3,4-thiadiazole and mixtures thereof; 2,5-bis(N,N-dialkyldithiocarbamoyl)-1,3,4-thiadiazoles; 2-alkyldithio-5-mercapto thiadiazoles; and the like. The metal deactivators may be used alone or in combination. The metal deactivators are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

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The anti-icing agents, includes but is not limited to di-ethylene glycol and the like. The anti-icing agents may be used alone or in combination. The anti-icing agents are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The demulsifiers, includes but is not limited to polyethylene and polypropylene oxide copolymers and the like. The demulsifiers may be used alone or in combination. The demulsifiers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The lubricity additives, includes but is not limited to glycerol monooleate, sorbitanmono oleate and the like. The lubricity additives may be used alone or in combination. The lubricity additives are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The friction modifiers, includes but is not limited to oleic acid and the like. The friction modifiers may be used alone or in combination. The friction modifiers are present in the range of about 0 % to about 90%, and in one embodiment about

.0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The flow improvers, includes but is not limited to ethylene vinyl acetate copolymers and the like. The flow improvers may be used alone or in combination. The flow improvers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025% to about 30% of the gel.

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The low temperature improvers, includes but is not limited to wax antisettling agents, ethylene vinyl acetate copolymers and the like. The low temperature improvers may be used alone or in combination. The low temperature improvers are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The cloud point depressants, includes but is not limited to alkylphenols and derivatives thereof, ethylene vinyl acetate copolymers and the like. The cloud point depressants may be used alone or in combination. The cloud point depressants are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The pour point depressant, includes but is not limited to alkylphenols and derivatives thereof, ethylene vinyl acetate copolymers and the like. The pour point depressant may be used alone or in combination. The pour point depressant are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The anti-static agents, includes but is not limited to polysiloxane polyether and the like. The anti-static agents may be used alone or in combination. The anti-static agents are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025 % to about 30% of the gel.

The valve seat recession agents, includes but is not limited to potassium or sodium bearing surfactants and the like. The valve seat recession agents may be used alone or in combination. The valve seat recession agents are present in the

range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025% to about 30% of the gel.

The intake valve deposit control additives, includes but is not limited to poly isobutylene amines and the like. The intake valve deposit control additives may be used alone or in combination. The intake valve deposit control additives are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025% to about 30% of the gel.

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The combustion chamber deposit control additives, includes but is not limited to polyetheramines and the like. The combustion chamber deposit control additives may be used alone or in combination. The combustion chamber deposit control additives are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025% to about 30% of the gel.

The fuel injector deposit control additives, includes but is not limited to alkylamines and the like. The fuel injector deposit control additives may be used alone or in combination. The fuel injector deposit control additives are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025% to about 30% of the gel.

The fuel dispersant additives, includes but is not limited to succinimides and the like. The fuel dispersant additives may be used alone or in combination. The fuel dispersant additives are present in the range of about 0 % to about 90%, and in one embodiment about .0005 % to about 50% and in another embodiment about .0025% to about 30% of the gel.

Optionally, an inert carrier can be used if desired. Furthermore, other active ingredients, which provide a beneficial and desired function to the soot being decreased, can also be included in the gel additive. In addition, solid, particulate additives such as the PTFE, MoS<sub>2</sub> and graphite can also be included.

The normally liquid hydrocarbon fuel may be a hydrocarbonaceous petroleum distillate fuel such as motor gasoline as defined by ASTM Specification D481 or diesel fuel or fuel oil as defined by ASTM Specification D975. Normally liquid hydrocarbon fuels comprising non-hydrocarbonaceous materials such as alcohols, ethers, organo-

nitro compounds and the like (e.g., methanol, ethanol, diethyl ether, methyl ethyl ether, methyl tert-butyl ether, nitromethane) are also include as are liquid fuels derived from vegetable or mineral sources such as corn, alfalfa, shale and coal. Normally liquid hydrocarbon fuels, which are mixtures of one or more hydrocarbonaceous fuels and one or more non-hydrocarbonaceous materials, are also included. Examples of such mixtures are combination of gasoline and ethanol, diesel fuel and ether, diesel fuel and methyl esters of vegetable or animal oils. In one embodiment, the fuel is a chlorine-free or low-chlorine fuel characterized by sulfur content of no more there about 10 ppm. Included are fuels known as gas to liquid fuels, GTL. The fuel may also be lead containing or lead free. The fuel may also be an emulsified fuel, either a macroemulsion, a micro-emulsion or combinations thereof.

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In an embodiment of this invention, the internal combustion engine is equipped with an exhaust after-treatment device. Exhaust after-treatment devices are used for modern engines to meet the new low exhaust emission standards. These systems are used to reduce undesirable emissions in the exhaust gases of internal combustion vehicle engines and are located in the exhaust system connected to the engines.

In one embodiment of this invention, catalysts are employed in the exhaust systems of internal combustion engines to convert carbon monoxide, hydrocarbons and nitrogen oxides (NOx) produced during engine operation into more desirable gases such as carbon dioxide, water and nitrogen. Among the broad range of available catalysts for this purpose, are oxidation catalysts, reduction catalysts and the so-called three-way converters. Oxidation catalysts can efficiently oxidize unburnt exhaust gas components and convert them into harmless substances. Three-way converters are able to simultaneously convert all three harmful substances provided that the internal combustion engine is operated close to the stoichiometirc air/fuel ratio. These catalyst systems typically contain noble metals from the platinum group of the Periodic System of Elements. Particular metals used are platinum, palladium and rhodium.

In another embodiment, the exhaust after-treatment device involves a NOx trap. NOx traps, i.e. materials that are able to absorb nitrogen oxides during lean-burn operation and are able to release them when the oxygen concentration in the exhaust gas

is lowered are porous support materials loaded with alkali metal or alkaline earth metals combined with precious metal catalysts such as platinum and the like.

In still another embodiment, the exhaust after-treatment device contains a diesel engine exhaust particulate filter hereinafter referred to as "DPF's". DPF's have a multiplicity of interconnected thin porous walls that define at least one inlet surface and one outlet surface on the filter and a multiplicity of hollow passages or cells extending through the filter from the inlet surface to an outlet surface. The interconnected thin porous walls allow the fluid to pass from the inlet surface to the outlet surface while restraining a desired portion of the solid particulates in the fluid from passing through. DPF's are typically installed in a housing which is inserted like a muffler or catalytic converter into the exhaust system of diesel engine equipped vehicle.

#### Specific Embodiment

In order to more thoroughly illustrate the present invention, the following examples are provided.

### 15 A. Gel Preparation

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A representative gel, known as Composition X is prepared by first mixing components A and C, and then adding component B with mixing in the proportions listed below. The resulting mixture is heated at 120° overnight to produce the final gel. The resulting gel is one of the formulations of the present invention.

20	Component	Chemical Description	% wt of Composition X
	Α	Polyisobutenyl (2000 Mn) succinimide Dispersant	20%
	В	400 TBN Overbased Alkylbenzenesulfonate Detergent	60%
25	C	Nonylated Diphenylamine Antioxidant	20%

From the above description and examples of the invention those skilled in the art will perceive improvements, changes and modifications in the invention. Such improvements, changes and modifications within the skill of the art are intended to be covered by the appended claims.

#### We claim:

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- 1. A composition comprising one or more fuel additives in a form of a gel used in an application selected from the group comprising decreasing the amount of soot in the lubricating oil engine, decreasing the amount of emissions in the engine exhaust and combinations thereof.
- 2. The composition of claim 1 wherein the gel comprises a dispersant, a detergent and an antioxidant.
- 3. The composition of claim 1 wherein the gel is represented by the formula A+B+C wherein A equals at least one component with at least one or more reactive or associative groups; wherein B equals a particle or other component with at least one group which reacts or associates with component A to form a gel and wherein C is at least one or more fuel additives.
- 4. The composition of claim 1 wherein the emissions reduced are selected from the group comprising soot, NOx, hydrocarbons and combinations thereof.
- 5. The composition of claim 3 wherein component A is selected from the group comprising antioxidants, dispersants, succinics, maleic anhydride styrene copolymers, maleated ethylene diene monomer copolymers, surfactants, emulsifiers, functionalized derivatives of such components and combinations thereof and in the range of about 0.1% to about 95% of the gel.
- 6. The composition of claim 3 wherein component B is selected from the group comprising dispersants, detergents, overbased detergents, carbon black, silica, alumina, titania, magnesium oxide, calcium carbonate, lime, clay, zeolites and combinations thereof and in the range of about 0.1 % of about 99% of the gel.
- 7. The composition of claim 3 wherein component C is selected from the group comprising antioxidants, extreme pressure agents, wear reduction agents, viscosity index improvers, anti-foaming agents, combustion modifiers and combinations thereof and in the range of about 0% to about 95% of the gel.
- 8. The composition of claim 7 further comprising at least one fuel additive selected from the group comprising friction reducing agents, extreme pressure (EP) agents, wear reduction agents, viscosity index improvers, anti-foaming agents, anti-

misting agents, cloud-point depressants, pour-point depressants, mineral or synthetic oils, anti-knock agents, lead scavengers, dyes, cetane improvers, rust inhibitors, bacteriostatic agents, gum inhibitors, fluidizers, metal deactivators, demulsifiers, anti-icing agents, lubricity additives, friction modifiers, viscosity improvers, flow improvers, low temperature improvers, anti-static agents, valve-seat recession agents, intake valve deposit control additives, combustion chamber deposit control additives, fuel injector deposit control additives and combinations thereof.

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- 9. The composition of claim 3 further comprising a fuel-borne catalysts (organometallic compounds of e.g. Na, K, Co, Ni, Fe, Cu, Mn, Mo, Va, Zi, Be, Pt, Pa, Ce, Cr, Al, Th, Se, Bi, Cd, Te, Th, Sn, Ba, B, La, Ta, Ti, W, Zn, Ga, Pb, Ag, Au, Os, Ir) and combinations thereof in the fuel.
- 10. The composition of claim 2 wherein the gel comprises an overbased detergent and an ashless succimide dispersant and wherein the ratio of detergent to dispersant is of about 10:1 to about 1:10 and wherein the total base number (TNB) of the overbased detergent is in the range from about 100 to about 400.
- 11. The composition of claim 2 when the dispersant is selected from the group comprising ashless succinimide, polyisobutenyl succinimide, substituted long chain alkenyl succinimides, high molecular weight esters, mannich dispersants, N-substituted long chain alkenyl succinimides, carboxylic dispersants, amine dispersants, polymeric dispersants, decyl methacrylate, vinyl decyl ether, aminoalkyl acrylates, acrylamides, poly-(oxyethylene)-substituted acrylates, high molecular weight olefins with monomers containing polar substitutes and a mixtures thereof; and a detergent selected from the group comprising overbased sulfonates, phenates, salicylates, carboxylates, overbased calcium sulfonate detergents, overbased detergents containing metals such as Mg, Ba, Sr, Na, Ca and K and mixtures thereof; and an antioxidant selected from the group comprises alkyl-substituted phenols, 2, 6-di-tertiary butyl-4-methyl phenol, phenate sulfides, phosphosulfurized terpenes, sulfurized esters, aromatic amines, hindered phenols and hindered, ester-substituted phenol and mixtures thereof.
- 12. A process comprising contacting in the range of a portion to all of the components of a gel of the composition of claim 3 in the combustion chamber of an

engine resulting in the reduction of soot in the engine oil, emissions in the engine exhaust and combinations thereof and wherein the emissions reduced in the exhaust are selected from the group comprising soot, NOx, hydrocarbons and combinations thereof.

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- 13. A process comprising contacting in the range of a portion to all of the components of a gel of the composition of claim 2 resulting in the reduction of soot in the engine oil and/or emissions in an engine exhaust.
- 14. The process of claim 12 wherein the gel is positioned to contact the fuel in an area selected from the group comprising full flow oil, bypass of oil, in the reservoir and combinations thereof.
- 15. The process of claim 12 wherein the gel is located in an area selected from the group comprising a filter, fuel filter, fuel bypass loop, fuel pump, injectors canister, housing, reservoir, pockets of a filter, canister in a filter, mesh in a filter, canister in a bypass system, mesh in a bypass system, canister in a tank, mesh in a tank, manifolds, inlets and/or outlets of fuel tank, fill fuel pipe, valves in fuel system, fuel chambers, fuel drain, intake air system, positive crank case ventilation system, air intake filter, exhaust gas recirculation (egr) system and combinations thereof.
- 16. The process of claim 15 wherein the gel is located in more than one
   location then the gel formulation can be identical, similar, different or combinations thereof.
  - 17. The process of claim 12 wherein the gel is in contact with the fuel system in the range of about 100% to 1% of the fuel in the engine.
  - 18. The process of claim 12 wherein the gel at the end of its service life contains in the range of none to a portion of the components in the gel remaining at the end of the service life of the gel due to selective dissolution of the gel.
  - 19. The process of claim 12 comprising adding the components of the gel to the fuel system in the range of all at the same time to a portion of the components of the gel over its service life.

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20. A fuel filter for an engine fuel system comprising a housing, a filter for removing particulate matter from a fuel filter and a container with a gel wherein the

gel comprises a dispersant, a detergent, and an antioxidant and results in the reduction of soot, emissions or combinations thereof from an engine.

- 21. A gel containment device for a fuel system comprising a housing in a fuel system and a container with a gel in the housing, and wherein the gel comprises a dispersant, a detergent and an antioxidant for the soot reduction, emissions reduction or combinations thereof from an engine.
  - 22. A fuel for an internal combustion engine comprising;
    - a) a fuel at 20°C and

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- b) a gel comprising a dispersant, a detergent and of an antioxidant, resulting in the soot reduction, emissions reduction or combinations thereof from the engine.
- 23. An internal combustion engine comprising
  - a) an internal combustion engine,
  - b) a liquid fuel at 20°C,
  - c) a gel comprising a dispersant, a detergent and of an antioxidant..
  - d) a reservoir containing at the gel supplying the gel component to the fuel in the fuel system of the engine resulting in the soot reduction, emissions reduction or combinations thereof from the engine.
- 24. An internal combustion engine according to claim 23, including an exhaust after treatment device that traps particulate (e.g. diesel particulate trap), oxidizes and/or reduces selected exhaust gas components, or traps or converts NOx to other compounds, or said engine is equipped with a system to recirculate exhaust gases to the intake air supply for said engine.
- 25. A fuel according to claim 22, wherein said fuel is characterized as a liquid fuel including diesel fuel, gasoline fuel, liquefied petroleum gas (LPG) an emulsified fuel, or combination thereof.
- 26. A fuel according to claim 22 wherein said fuel is characterized as a gaseous fuel, including natural gas, methane, ethane, propane or combinations thereof.

## INTERNATIONAL SEARCH REPORT

Intern 1al Application No PC I / US 2004/019819

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C10L10/02 C10L10/06 C10L1/12 C10L1/14 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) IPC 7 C10L Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with Indication, where appropriate, of the relevant passages Relevant to claim No. WO 2004/072214 A (LUBRIZOL CORP; LAWATE E 1-8 SAURABH S (US); SILVERSTEIN ROBERT (US))
26 August 2004 (2004-08-26) page 4, line 28 - line 30; claims 1,12,32,33 P,X WO 2004/007653 A (LUBRIZOL CORP) 22 January 2004 (2004-01-22) 1-11 claims 1-7; table 1 Χ US 4 382 005 A (MILLER HAROLD N) 1-11,22,3 May 1983 (1983-05-03) column 7, line 15 - line 43 column 8, line 32 - line 46; claim 1 25,26 Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "O" document referring to an oral disclosure, use, exhibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report . 22 H. C. 23 September 2004 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Bertrand, S

## INTERNATIONAL SEARCH REPORT

al Application No PC1, u32004/019819

	ation) DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category °	Citation of document, with indication, where appropriate, of the relevant passages	nelevant to daim No.
Χ .	US 3 342 733 A (ROBBINS MAX L ET AL) 19 September 1967 (1967-09-19)	1,3-9, 11,22, 25,26
i	column 2, line 28 - line 39 column 3, line 50 - line 70 column 7, line 58 - line 61	
Х	EP 0 476 197 A (ETHYL PETROLEUM ADDITIVES LTD) 25 March 1992 (1992-03-25)	12-14, 17-19, 22,25,26
	page 3, line 57 - page 10, line 47; claim 1	
X	EP 0 423 744 A (LUBRIZOL CORP) 24 April 1991 (1991-04-24)	12-14, 17-19, 22,25,26
	page 23, line 9 - line 22; claims 1,5,9,18	12,20,20
X	DATABASE WPI Section Ch, Week 198850 Derwent Publications Ltd., London, GB; Class A97, AN 1988-358626 XP002297647	1,3-8
	& JP 63 270797 A (MITSUBISHI OIL CO) 8 November 1988 (1988-11-08) abstract	
X	EP 0 516 461 A (ETHYL PETROLEUM ADDITIVES INC) 2 December 1992 (1992-12-02) example XX	1-11
Χ .	EP 0 277 729 A (AMOCO CORP) 10 August 1988 (1988-08-10) example 1; table 1	1-11
	·	

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## INTERNATIONAL SEARCH REPORT

ational application No. PCT/US2004/019819

Box II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)
This international Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
Claims Nos.:     because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:  1-19, 22, 25, 26
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 1-19,22,25,26

Gel composition of claim 1. process for reducing soot and emisssions by using the gel composition. Fuel composition comprising the components of the gel composition.

2. claims: 20, 21

Fuel filter of claim 20. Gel containment device of claim 21.

3. claims: 23,24

Internal combustion engine of claim 23

# INTERNATIONAL SEARCH REPORT remails on on patent family members

Intern al Application No PC1/uS2004/019819

atent document	t t				2004/019819
d in search report		Publication date		Patent family member(s)	Publication date
2004072214	. A	26-08-2004	WO	2004072214 A1	26-08-2004
2004007653	A 	22-01-2004	US WO	2004014614 A1 2004007653 A2	22-01-2004 22-01-2004
4382005	A .	03-05-1983	AU AU CA DE EP JP JP	547946 B2 7619781 A 1180850 A1 3164748 D1 0049975 A2 1667822 C 3032597 B	14-11-1985 22-04-1982 08-01-1985 16-08-1984 21-04-1982 29-05-1992
			JP	57092091 A	13-05-1991 08-06-1982
3342733	Α	19-09-1967	NONE		
0476197	Α	25-03-1992	EP AU AU CA DE DE ES JP JP	0476197 A1 634422 B2 8465391 A 2051451 A1 69006029 D1 69006029 T2 2048439 T3 2965764 B2 4234490 A	25-03-1992 18-02-1993 26-03-1992 21-03-1992 24-02-1994 05-05-1994 16-03-1994 18-10-1999 24-08-1992
0423744	A	24-04-1991	USTPRTTUUURA AAA AAB CCD DE DE DE DE DE DE EEFFIKLP.	4690687 A 4659338 A 77828 T 0423744 A1 243589 A1 118528 T 154068 T 591394 B2 6192986 A 600058 B2 8253087 A 8606850 A 1303853 C 86106817 A ,B 3650239 D1 3650239 T2 3650634 D1 3650634 T2 3685877 D1 3685877 T3 233250 T1 66695 A 192587 A 0233250 A1 0579339 A1 2001515 A6 871661 A 871707 A 36993 A 79599 A 7088514 B	01-09-1987 21-04-1987 15-07-1992 24-04-1991 31-08-1993 15-03-1995 15-06-1997 30-11-1989 10-03-1987 02-08-1990 31-03-1988 03-11-1987 23-06-1992 27-05-1987 23-03-1995 08-06-1995 10-07-1997 15-01-1998 06-08-1992 29-07-1999 17-03-1988 12-06-1995 16-04-1987 26-08-1987 19-01-1994 01-06-1988 15-04-1987 23-04-1987 16-04-1987 23-04-1993 31-01-1991 27-09-1995 03-03-1988
•	9423744	0423744 A	9423744 A 24-04-1991	0423744 A 24-04-1991 US US AT EP AR AT AT AU AU AU AU BR CA CN DE DE DE DE DE DE DE DE DE FI FI HK IL JP	0423744 A 24-04-1991 US 4690687 A US 4659338 A AT 77828 T EP 0423744 A1 AR 243589 A1 AT 118528 T AT 118528 T AT 154068 T AU 591394 B2 AU 6192986 A AU 600058 B2 AU 6192986 A CA 1303853 C CN 86106817 A BR 8606850 A CA 1303853 C CN 86106817 A BDE 3650239 D1 DE 3650634 D1 DE 3650634 D1 DE 3650634 D1 DE 3685877 D1 D1 DE 3685877 D1

# INTERNATIONAL SEARCH REPORT mation on patent family members

Intern al Application No PCT/US2004/019819

Patent document cited in search report	Publication date		Patent family member(s)	Publication date ·
EP 0423744 A		MX NO NO SG WO IN US ZA	164983 B 871551 A ,B, 952019 A 15993 G 8701126 Al 167837 Al 4804389 A 8606177 A	13-10-1992 13-04-1987 22-05-1995 16-04-1993 26-02-1987 29-12-1990 14-02-1989 25-03-1987
JP 63270797 A	08-11-1988	JP · JP	1904925 C 6033385 B	08-02-1995 02-05-1994
EP 0516461 A	02-12-1992	AU CA DE DE EP JP	657528 B2 1523492 A 2065945 A1 69208526 D1 69208526 T2 0516461 A1 5179273 A	16-03-1995 03-12-1992 30-11-1992 04-04-1996 13-06-1996 02-12-1992 20-07-1993
EP 0277729 A	10-08-1988	DE EP	3868949 D1 0277729 A1	16-04-1992 10-08-1988

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